510(K) SUMMARY OF SAFETY AND EFFECTIVENESS

This summary of safety and effectiveness is provided as part of this Premarket Notification in compliance with 21 CFR, Part 807, Subpart E, Section 807.92.

1. Submitter's Information: 21 CFR 807.92(a)(1)

MEDISON CO., LTD. 1003, Daechi-dong, Gangnam-gu, Seoul 135-280; Korea

AUG 0 6 2010

Contact Person:

Kyeong-Mi, Park Regulatory Affairs Manager

Telephone:

82.2.2194.1373

Facsimile:

82.2.556.9209

Data Prepared: July 2, 2010

2. Name of the device:

Common/Usual Name:

Diagnostic Ultrasound System and Accessories

Proprietary Name:

SONOACE R7 Diagnostic Ultrasound System

Classification Names:	FR Number	Product Code
Ultrasonic Pulsed Doppler Imaging System	892.1550	IYN
Ultrasound Pulsed Echo Imaging System	892.1560	IYO
Diagnostic Ultrasound Transducer	892.1570	ITX

3. Identification of the predicate or legally marketed device:

K100186, MySono U5 Diagnostic Ultrasound System K093714, SONOACE X8 Diagnostic Ultrasound System

4. Device Description:

The SONOACE R7 is a general purpose, mobile, software controlled, diagnostic ultrasound system. Its function is to acquire ultrasound data and to display the data as B mode, M mode, Color Doppler imaging, Power Doppler imaging, PW/CW Spectral Doppler mode, Harmonic imaging, 3D imaging mode or as a combination of these modes. The SONOACE R7 also gives the operator the ability to measure anatomical structures and offers analysis packages that provide information that is used to make a diagnosis by competent health care professionals. The SONOACE R7 has real time acoustic output display with two basic indices, a mechanical index and a thermal index, which are both automatically displayed.

The SONOACE R7 has been designed to meet the following product safety standards:

- UL 60601-1, Safety requirements for Medical Equipment
- CSA C22.2 No. 601.1, Safety requirements for Medical Equipment
- IEC60601-2-37, Diagnostic Ultrasound Safety Standards
- EN/IEC60601-1, Safety requirements for Medical Equipment
- EN/IEC60601-1-2, EMC requirements for Medical Equipment
- NEMA UD-2, Acoustic Output Measurement Standard for Diagnostic Ultrasound Equipment
- NEMA UD-3, Standard for Real Time Display of Thermal and Mechanical Acoustic Output Indices on Diagnostic Ultrasound Equipment
- IEC 61157, Declaration of the acoustic output
- ISO10993-1, Biocompatibility

5. Intended Uses:

1

The SONOACE R7 Diagnostic Ultrasound System and transducers are intended for diagnostic ultrasound imaging and fluid analysis of the human body.

The clinical applications include: Fetal, Abdominal, Pediatric, Small Organs, Adult Cephalic, Transrectal, Trans-vaginal, Muscular-Skeletal (Conventional, Superficial), Cardiac Adult, Cardiac Pediatric, Peripheral vessel.

6. Technological Characteristics:

The SONOACE R7 is substantially equivalent to the SONOACE X8 Diagnostic Ultrasound System, cleared via K093714, and the MySono U5 Diagnostic Ultrasound System, cleared via K100186. All systems transmit ultrasonic energy into patients, then perform post processing of received echoes to generate on-screen display of anatomic structures and fluid flow within the body. All system allow for specialized measurements of structures and flow, and calculations.

END of 510(K) Summary

DEPARTMENT OF HEALTH & HUMAN SERVICES



Food and Drug Administration 10903 New Hampshire Avenue Document Control Room --WO66-G609 Silver Spring, MD 20993-0002

Medison Co., Ltd. % Mr. Mark Job Responsible Third Party Official Regulatory Technology Services LLC 1394 25th Street BUFFALO MN 55313

AUG 0 6 2010

Re: K102065

Trade/Device Name: SONOACE R7 Diagnostic Ultrasound System

Regulation Number: 21 CFR 892.1550

Regulation Name: Ultrasonic pulsed doppler imaging system

Regulatory Class: II

Product Code: IYN, IYO, and ITX

Dated: July 22, 2010 Received: July 23, 2010

Dear Mr. Job:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and we have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration.

This determination of substantial equivalence applies to the following transducers intended for use with the SONOACE R7 Diagnostic Ultrasound System, as described in your premarket notification:

Transducer Model Number

C2-8 ER4-9/10ED EV4-9/10ED L3-8 L5-12/50EP HL5-12ED P2-4AH 3DC2-6 3D4-8ET

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to such additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 895. In addition, FDA may publish further announcements concerning your device in the Federal Register.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

This letter will allow you to begin marketing your device as described in your premarket notification. The FDA finding of substantial equivalence of your device to a legally marketed predicate device results in a classification for your device and thus permits your device to proceed to market.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please go to http://www.fda.gov/AboutFDA/CentersOffices/CDRH/CDRHOffices/ucm115809.htm for the Center for Devices and Radiological Health's (CDRH's) Office of Compliance. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to

http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

If you have any questions regarding the content of this letter, please contact Jana Delfino at (301) 796-6503.

Sincerely yours.

Donald J. St.Pierre

Acting Director

Division of Radiological Devices Office of *In Vitro* Diagnostic Device

Evaluation and Safety

Center for Devices and Radiological Health

Enclosure(s)

SECTION 1.3 INDICATIONS FOR USE

510(k) Number	· (if known): _	K1020	065	. A	UG 0 6 2010
Device Name:	SONOACE	R7 Diagnostic U	Itrasound System		
Indications for	Use:	-	•		
and fluid analys The clinical ap	sis of the hum plications incl	an body. ude: Fetal, Abdor	minal, Pediatric, Sma	are intended for diagnostic all Organ, Adult Cephalic, T dult, Cardiac Pediatric, Peri	rans-rectal. Trans-
Prescripti (Part 21 C	on Use	art D)	AND/OR	Over-The-Counte (21 CFR 801 Sub	
, (PLE	ASE DO NOT	WRITE BELOW	THIS LINE-CONTINU	JE ON ANOTHER PAGE OF	NEEDED)
WESTERS SECTIONS OF THE SECTION OF T	. Concu	rence of CDRH,	Office of In Vitro D	iagnostic Devices (OIVD)	
Indications for Use	e	. Office of	(Division Sign Division of Radiologi of In Vitro Diagnostic Devic	ical Devices ce Evaluation and Safety	Section 1.3, page 1

510(k) No.:

Device Name: SONOACE R7 Diagnostic Ultrasound System

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Clinical Application					Operation (*inc	cludes simultaneou	
General (Track Lonly)	Specific (Tracks I & III)	В	М	PWD	CWD	Color Doppler*	Combined* (Spec.)	Other (Spec.)
Ophthalmic	Ophthalmic							
	Fetal (See Note 3)	N	N	N		N	Note I	Notes 2, 7, 8
	Abdominal	Ν	N	N	N	N	Note I	Notes 2, 4, 7, 8
	Intra-operative (See Note 6)							
	Intra-operative (Neuro.)	T						
Fetal Imaging	Laparoscopie							
& Other	Pediatric	N	N	N		. N	Note 1	Note 2, 5, 6, 7, 8, 9
	Small Organ (See Note 5)	N	N	N		N	Note I	Note 2, 5, 6, 7, 9
	Neonatal Cephalic							
	Adult Cephalic	N	N	N	N	N	Note I	Note 4, 7
	Trans-rectal	N	N	N		N	Note 1	Note 2, 8
	Trans-vaginal	N	N	N		N	Note I	Note 2, 8
	Trans-urethral							
	Trans-esoph. (non-Cardiac)	7						
	Musculo-skel. (Convent.)	N	N	N		N	Note I	Note 2, 5, 6, 7, 9
	Musculo-skel. (Superfic.)	N	N	. N		N	Note I	Note 2, 5, 6, 7, 9
	Intra-luminal							
	Other (spec.)							
	Cardiac Adult	N	N	N	N	N	Note 1	Note 4, 7
Cardiac	Cardiac Pediatric	N	И	N	N	N	Note I	Note 4, 7
	Trans-esophageal (Cardiac)			L				
	Other (spec.)							
Peripheral	Peripheral vessel	N	N	N		N	Note 1	Note 2, 5, 6, 7, 9
Vessel	Other (spec.)							

N= new indication; P= previously cleared by FDA; E= added under Appendix E

Additional Comments:

Color Doppler includes Power (Amplitude) Doppler

Note I: B/M, B/PWD, B/Color Doppler, B/Color Doppler/PWD, B/Power Doppler/PWD, B/Color Doppler/Color M

Note 2: Includes imaging for guidance of biopsy

Note 3: Includes infertility monitoring of follicle development

Note 4: Color M-mode

Note 5: For example, thyroid, parathyroid, breast, scrotum and penis in adult, pediatric and neonatal patients

Note 6: Abdominal organs and peripheral vessel

Note 7: Tissue Harmonic Imaging (FIII)

Note 8: 3D imaging

Note 9: Panoramic imaging

Concurrence of CDRH, Office of In Vitro Diagnostic Devices (OIVD) Prescription Use (Per 21 CFR 801.109)

Indications for Use

Section 1.3, page 2

510(k) No.:

Device Name: C2-8 for use with SONOACE R7

Intended Lice. Diagnostic ultracound imaging or fluid flow analysis of the human hody as follows:

	Diagnostic ultrasound in Clinical Application	laging	OI I	iuid ne			des simultaneous B	
General (Track Lonly)	Specific (Tracks I & III)	В	М	PWD	CWD	Color Doppler*	Combined* (Spec.)	Other (Spec.)
Ophthalmic	Ophthalmic							
	Fetal (See Note 3)	Ь	Р	Р		Р	Note I	Notes 2, 7, 8
	Abdominal .	Р	þ	Р		P	Note I	Notes 2, 7, 8
	Intra-operative (See Note 6)							
	Intra-operative (Neuro.)							
Fetal Imaging	Laparoscopie					1		
& Other	Pediatric	Р	Р	l ₃		Р.	Note I	Notes 2, 7, 8
	Small Organ (See Note 5)		i					
	Neonatal Cephalic							
	Adult Cephalic	$\neg \vdash$		·				
	Trans-rectal	_						
	Trans-vaginal	\top		1				
	Trans-urethral			<u> </u>				· ······· ,
	Trans-esoph. (non-Cardiac)	\top		<u> </u>				
	Musculo-skel. (Convent.)	\top						
	Musculo-skel. (Superfic.)	\neg			· · · · · ·		1	
	Intra-luminal		-	<u> </u>				
	Other (spec.)						ii	···· · · ·
-	Cardiac Adult			1			1	
Cardiac	Cardiac Pediatric							
	Trans-esophageal (Cardiac)	T			<u> </u>			
	Other (spec.)				ļ			
Peripheral	Peripheral vessel							
Vessel	Other (spec.)		Γ				1	

N= new indication; P= previously cleared by FDA K093714; E= added under Appendix E

Additional Comments:

Color Doppler includes Power (Amplitude) Doppler

Note 1: B/M, B/PWD, B/Color Doppler, B/Color Doppler/PWD, B/Power Doppler/PWD, B/Color Doppler/Color M

Note 2: Includes imaging for guidance of biopsy Note 3: Includes infertility monitoring of follicle development

Note 4: Color M-mode

Note 5: For example: thyroid, parathyroid, breast, scrotum and penis in adult, pediatric and neonatal patients Note 6: Abdominal organs and peripheral vessel Note 7: Tissue Harmonic Imaging (THI)

Note 8: 3D imaging Note 9: Panoramic imaging

Concurrence of CDRH, Office of In Vitro Diagnostic Devices (OIVD) Prescription Use (Per 21 CFR 801.109)

Indications for Use

Section 1.3, page 3

510(k) No.:

Device Name: ER4-9/10ED for use with SONOACE R7

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Clinical Application						des simultaneous B-	mode)
General (Track Lonly)	Specific (Tracks I & III)	В	М	PWD	CWD	Cotor Doppler*	Combined* (Spec.)	Other (Spec.)
Ophthalmic	Ophthalmic							
	Fetal (See Note 3)				·		-	
	Abdominal							
	Intra-operative (See Note 6)							
	Intra-operative (Neuro.)							
Fetal Imaging	Laparoscopic		<u> </u>					
& Other	Pediatric			i –				
	Small Organ (See Note 5)	1						
	Neonatal Cephalic							
	Adult Cephalic			!				
	Trans-rectal	P	Р	P		Р	Note 1	Notes 2, 8
	Trans-vaginal	Р	Р	Р		р	Note 1	Notes 2, 8
	Trans-urethral			 	·			
	Trans-esoph, (non-Cardiac)	\neg		—				
	Musculo-skel, (Convent.)							•
	Musculo-skel. (Superfic.)			 -		<u> </u>	<u> </u>	
	Intra-luminal			i i			i	· ···· · · · ·
•	Other (spec.)						1	
	Cardiac Adult							
Cardiac	Cardiac Pediatric				-			
	Trans-esophageal (Cardiac)			1				
	Other (spec.)			<u> </u>			1	
Peripheral	Peripheral vessel							
Vessel	Other (spec.)	1				<u> </u>		

N= new indication; P= previously cleared by FDA K093714; E= added under Appendix E

Additional Comments:

Color Doppler includes Power (Amplitude) Doppler

Note 1: B/M, B/PWD, B/Color Doppler, B/Color Doppler/PWD, B/Power Doppler/PWD, B/Color Doppler/Color M

Note 2: Includes imaging for guidance of biopsy
Note 3: Includes infertility monitoring of follicle development

Note 4: Color M-mode

Note 5: For example: thyroid, parathyroid, breast, scrotum and penis in adult, pediatric and neonatal patients
Note 6: Abdominal organs and peripheral vessel
Note 7: Tissue Harmonic Imaging (TIII)

Note 8: 3D imaging

Note 9: Panoramic imaging

Concurrence of CDRH, Office of In Vitro Diagnostic Devices (OIVD) Prescription Use (Per 21 CFR 801.109)

Indications for Use

(Division Sign-Off) Division of Radiological Devices Office of In Vitro Diagnostic Device Evaluation and Safety

Section 1.3, page 4

510(k) No.:

Device Name: EV4-9/10ED for use with SONOACE R7

Intended Use: Diagnostic ultrasound imaging or fluid flow

	Clinical Application				Mode of C	peration (*inclu	des simultaneous B	-mode)
General (Track Lonly)	Specific (Tracks I & III)	В	М	PWD	CMĐ	Color Doppler*	Combined* (Spec.)	Other (Spec.)
Oplithalmic	Ophthalmic							
	Fetal (See Note 3)							
	Abdominal							
	Intra-operative (Sec Note 6)							· · · · · · · · · · · · · · · · · · ·
	Intra-operative (Neuro.)							
Fetal Imaging	Laparoscopic		·					
& Other	Pediatric							
	Small Organ (See Note 5)					1		
•	Neonatal Cephalic	ı						
	Adult Cephalic							
	Trans-rectal	P	P	Р		Р	Note I	Notes 2, 8
	Trans-vaginal	P	p	Р		Р	Note I	Notes 2, 8
	Trans-urethral	T						
	Trans-esoph. (non-Cardiac)					<u> </u>		
•	Musculu-skel, (Convent.)				<u> </u>			
	Musculo-skel. (Superfic.)			1				
	Intra-luminal	Ti -						
	Other (spec.)							
	Cardiac Adult				Ĭ			•
Cardiac	Cardiac Pediatric				L			_
	Trans-esophageal (Cardiac)							
	Other (spec.)							
Peripheral	Peripheral vessel							
Vessel	Other (spec.)	1						

N= new indication; P= previously cleared by FDA K093714; E= added under Appendix E

Additional Comments:

Color Doppler includes Power (Amplitude) Doppler

- Note 1: B/M, B/PWD, B/Color Doppler, B/Color Doppler/PWD, B/Power Doppler/PWD, B/Color Doppler/Color M
- Note 2: Includes imaging for guidance of biopsy
- Note 3: Includes infertility monitoring of follicle development
- Note 4: Color M-mode
- Note 5: For example: thyroid, parathyroid, breast, scrotum and penis in adult, pediatric and neonatal patients
- Note 6: Abdominal organs and peripheral vessel Note 7: Tissue Harmonic Imaging (TIII) Note 8: 3D imaging

- Note 9: Panoramic imaging

Concurrence of CDRH, Office of In Vitro Diagnostic Devices (OIVD) Prescription Use (Per 21 CFR 801.109)

Indications for Use

Section 1.3, page 5

(Division Sign-Off) Division of Radiological Devices Office of In Vitro Diagnostic Device Evaluation and Safety

510K

510(k) No.:

Device Name: L3-8 for use with SONOACE R7

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Clinical Application				Mode of C	peration (*inclu	des simultaneous B	-mode)
General (Track Lonly)	Specific (Tracks I & HI)	В	М	рwb	CMD	Color Doppler*	Combined* (Spec.)	Other (Spec.)
Ophthalmic	Ophthalmic						,	
	Fetal (See Note 3)							
	Abdominat							-
	Intra-operative (See Note 6)							
	Intra-operative (Neuro.)							
Fetal Imaging	Laparoscopie							
& Other	Pediatric	N	N	N		N	Note I	Note 2, 5, 6, 7, 9
	Small Organ (See Note 5)	N	И	N		N	Note I	Note 2, 5, 6, 7, 9
	Neonatal Cephatic							
	Adult Cephalic							
	Trans-rectal							
	Trans-vaginal							
	Trans-urethral							·· , . <u></u> .
	Trans-esoph, (non-Cardiac)							
	Musculo-skel, (Convent.)	N	N	N	1	N	Note I	Note 2, 5, 6, 7, 9
	Musculo-skel, (Superfic.)	N	N	N		N	Note I	Note 2, 5, 6, 7, 9
	Intra-luminal							
	Other (spec.)							
	Cardiac Adult							
Cardiac	Cardiac Pediatric				<u> </u>			
	Trans-esophageal (Cardiac)							
	Other (spec.)				İ			
Peripheral	Peripheral vessel	N	N	N		N	Note I	Note 2, 5, 6, 7, 9
Vessel	Other (spec.)							

N= new indication; P= previously cleared by FDA; E= added under Appendix E

Additional Comments:

Color Doppler includes Power (Amplitude) Doppler

Note 1: B/M, B/PWD, B/Color Doppler, B/Color Doppler/PWD, B/Power Doppler/PWD, B/Color Doppler/Color M

Note 2: Includes imaging for guidance of biopsy

Note 3: Includes infertility monitoring of follicle development

Note 4: Color M-mode

Note 5: For example: thyroid, parathyroid, breast, scrotum and penis in adult, pediatric and neonatal patients

Note 6: Abdominal organs and peripheral vessel

Note 7: Tissue Harmonic Imaging (THI)

Note 8: 3D imaging

Note 9: Panoramic imaging

Concurrence of CDRH, Office of In Vitro Diagnostic Devices (OIVD)
Prescription Use (Per 21 CFR 801.109)

Indications for Use

(Division Sign-Off)

Division of Radiological Devices

Office of In Vitro Diagnostic Device Evaluation and Safety

510K K102065

Section 1.3, page 6

510(k) No.:

Device Name: L5-12/50EP for use with SONOACE R7

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Clinical Application				Mode of C	peration (* inclu	des simultaneous B	-mode)
General (Track Lonly)	Specific (Tracks I & III)	В	М	טשין .	CWD	Color Doppler*	Combined* (Spec.)	Other (Spec.)
Ophthalmic	Ophthalmic							··-
**-	Fetal (See Note 3)	1						
	Abdominal							
	Intra-operative (See Note 6)							
	Intra-operative (Neuro.)							
etal Imaging	Laparoscopic		i —				-	· · · · ·
& Other	Pediatric	P	Р	Ь		Р	Note 1	Note 2, 5, 6, 7, 9
	Small Organ (See Note 5)	Р	p	Р		Ρ.	Note 1	Note 2, 5, 6, 7, 9
	Neonatal Cephalic		1					
	Adult Cephalic	1						
	Trans-rectal		<u> </u>					
	Trans-vaginal			· ·				
	Trans-urethral		<u> </u>	1				
	Trans-esoph. (non-Cardiac)	_			_			
	Musculo-skel. (Convent.)	Р	Р	Р		Р	Note I	Note 2, 5, 6, 7, 9
	Musculo-skel. (Superfic.)	Р	Р	Р		Р	Note 1	Note 2, 5, 6, 7, 9
	Intra-luminal			i				
	Other (spec.)							
	Cardiac Adult							
Cardiac	Cardiac Pediatric							·
	Trans-esophageal (Cardiac)							
	Other (spec.)							
Peripheral	Peripheral vessel	P	P	Р		Р	Note I	Note 2, 5, 6, 7, 9
Vessel	Other (spec,)							

N= new indication; P= previously cleared by FDA K093714; E= added under Appendix E

Additional Comments:

Color Doppler includes Power (Amplitude) Doppler

Note 1: B/M, B/PWD, B/Color Doppler, B/Color Doppler/PWD, B/Power Doppler/PWD, B/Color Doppler/Color M

Note 2: Includes imaging for guidance of biopsy

Note 3: Includes infertility monitoring of follicle development

Note 4: Color M-mode

Note 5: For example: thyroid, parathyroid, breast, scrotum and penis in adult, pediatric and neonatal patients

Note 6: Abdominal organs and peripheral vessel

Note 7: Tissue Harmonic Imaging (THI)

*Note 8: 3D imaging

Note 9: Panoramic imaging

Concurrence of CDRH, Office of In Vitro Diagnostic Devices (O1VD)
: Prescription Use (Per 21 CFR 801.109)

Indications for Use

Section 1.3, page 7

(Division Sign-Off)
Division of Radiological Devices
Office of In Vitro Diagnostic Device Evaluation and Safety

610K 5102065

510(k) No.:

Device Name: HL5-12ED for use with SONOACE R7

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Clinical Application						des simultaneous B	-mode)
General (Track Lonly)	Specific (Tracks I & III)	В	М	PWD	CWD	Cotor Doppler*	Combined* (Spec.)	Other (Spec.)
Opluhalmic	Ophthalmic							•
	Fetal (See Note 3)							
	Abdominal							
	Intra-operative (See Note 6)							
	Intra-operative (Neuro.)							
fetal Imaging	Laparoscopic		_			· · · · · · · · · · · · · · · · · · ·		
& Other	Pediatric	Р	P	Р		Р	Note 1	Note 2, 5, 6, 7, 9
	Small Organ (See Note 5)	Р	Р	Р		р	Note 1	Note 2, 5, 6, 7, 9
	Neonatal Cephalic							
	Adult Cephalic							
	Trans-rectal							,,
	Trans-vaginal			1	-			
	Trans-urethral							
	Trans-esoph (non-Cardiac)			T				
	Musculo-skel, (Convent.)	Р	Р	P		Р	Note I	Note 2, 5, 6, 7, 9
	Musculo-skel. (Superfic.)	þ	Р	Р		Ρ	Note I	Note 2, 5, 6, 7, 9
	Intra-luminal		İ					
	Other (spec.)	1	\Box					
	Cardiac Adult							
Cardiac	Cardiac Pediatric							
	Trans-esophageal (Cardiac)							
	Other (spec.)							
Peripheral	Peripheral vessel	þ	Р	Ρ		Р	Note 1	Note 2, 5, 6, 7, 9
Vessel	Other (spec.)							

N= new indication; P= previously cleared by FDA K093714; E= added under Appendix E

Additional Comments:

Color Doppler includes Power (Amplitude) Doppler

Note 1: B/M, B/PWD, B/Color Doppler, B/Color Doppler/PWD, B/Power Doppler/PWD, B/Color Doppler/Color M

Note 2: Includes imaging for guidance of biopsy

Note 3: Includes infertility monitoring of follicle development

Note 4: Color M-mode

Note 5: For example: thyroid, parathyroid, breast, scrotum and penis in adult, pediatric and neonatal patients

Note 6: Abdominal organs and peripheral vessel

Note 7: Tissue Harmonic Imaging (THI)

Note 8: 3D imaging

Note 9: Panoramic imaging

Concurrence of CDRH, Office of In Vitro Diagnostic Devices (OIVD)
Prescription Use (Per 21 CFR 801.109)

Indications for Use

Section 1.3, page 8

(Division Sign-Off)
Division of Radiological Devices
Office of In Vitro Diagnostic Device Evaluation and Safety

610K 15102065

510(k) No.:

Device Name: P2-4AH for use with SONOACE R7

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Clinical Application				Mode of C	Operation (*inclu	des simultaneous B-r	
General (Track Lonly)	Specific (Tracks I & III)	į,	М	PWD	CWD	Color Doppler*	Combined* (Spec.)	Other (Spec.)
Ophthalmic	Ophthalmic							
	Fetal (See Note 3)							
	Abdominat	, P	Р	P	Р	Р	Note I	Note 4, 7
	Intra-operative (See Note 6)							· <u></u> ·
	Intra-operative (Neuro.)							
Fetal Imaging	Laparoscopic							
& Other	Pediatric							•
	Small Organ (See Note 5)	`` `						
	Neonatal Cephalic	\top						
	Adult Cephalic	Р	Р	P	p	Р	Note I	Note 4, 7
	Trans-rectal		_					
	Trans-vaginal					· · ·		
	Trans-urethral							
	Trans-esoph (non-Cardiac)	1			1	1		
	Musculo-skel. (Convent.)							
•	Musculo-skel. (Superfic.)					,		-
	Intra-luminal			_	 			
	Other (spec.)							
	Cardiac Adult	Р	Р	р	Р	Р	Note I	Note 4, 7
Cardiac	Cardiac Pediatric	Р	Р	Р	P	Р	Note 1	Note 4, 7
	Trans-esophageal (Cardiac)							
	Other (spec.)							
Peripheral	Peripheral vessel							
Vessel	Other (spec.)							

N= new indication; P= previously cleared by FDA K093714; E= added under Appendix E

Additional Comments:

Color Doppler includes Power (Amplitude) Doppler

Note 1: B/M, B/PWD, B/Color Doppler, B/Color Doppler/PWD, B/Power Doppler/PWD, B/Color Doppler/Color M

Note 2: Includes imaging for guidance of biopsy

Note 3: Includes infertility monitoring of follicle development

Note 4: Color M-mode

Note 5: For example: thyroid, parathyroid, breast, scrotum and penis in adult, pediatric and neonatal patients

Note 6: Abdominal organs and peripheral vessel

Note 7: Tissue Harmonic Imaging (TIII)

Note 8: 3D imaging

Note 9: Panoramic imaging

Concurrence of CDRH, Office of In Vitro Diagnostic Devices (OIVD)
Prescription Use (Per 21 CFR 801.109)

Indications for Use

Section 1.3, page 9

(Division Sign-Off)
Division of Radiological Devices
Office of In Vitro Diagnostic Device Evaluation and Safety

510K KLO20G5

510(k) No.:

Device Name: 3DC2-6 for use with SONOACE R7

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis of the human body as follows:

	Clinical Application						des simultaneous B-	
General (Track Lonly)	Specific (Tracks I & III)	В	М	PWD	CWD	Color Doppler*	Combined* (Spec.)	Other (Spec.)
Ophthalmic	Ophthalmic					•		
	Fetal (See Note 3)	þ	Р	P		þ	Note 1	Note 2, 7, 8
	Abdominal	P	P	p.		P	Note 1	Note 2, 7, 8
	Intra-operative (Nee Note 6)							
	Intra-operative (Neuro.)							
Fetal Imaging	Laparoscopic							
& Other	Pediatric	P	P	Ь		P	Note I	Note 2, 7, 8
	Small Organ (See Note 5)			1				
	Neonatal Cephalic			1		-		
	Adult Cephalic		1					•
•	Trans-rectal							
	Trans-vaginal							
	Trans-urethral			1		1		
	Trans-esoph. (non-Cardiac)							
	Musculo-skel. (Convent.)	1	1	 	_		1	
	Musculo-skel. (Superfic.)	_	1					
	Intra-luminal							
	Other (spec.)							
<u></u>	Cardiae Adult							
Cardiac	Cardiac Pediatric	1		`	i			
	Trans-esophageal (Cardiac)	1			1			
	Other (spec.)						<u> </u>	
Peripheral	Peripheral vessel							
Vessel	Other (spec.)		1			_		

N= new indication; P= previously cleared by FDA K093714; E= added under Appendix E

Additional Comments:

Color Doppler includes Power (Amplitude) Doppler

Note 1: B/M, B/PWD, B/Color Doppler, B/Color Doppler/PWD, B/Power Doppler/PWD, B/Color Doppler/Color M

Note 2: Includes imaging for guidance of biopsy Note 3: Includes infertility monitoring of follicle development

Note 4: Color M-mode

Note 5: For example: thyroid, parathyroid, breast, scrotum and penis in adult, pediatric and neonatal patients

Note 6: Abdominal organs and peripheral vessel

Note 7: Tissue Harmonic Imaging (TIII) Note 8: 3D imaging

Note 9: Panoramic imaging

Concurrence of CDRH, Office of In Vitro Diagnostic Devices (OIVD) Prescription Use (Per 21 CFR 801.109)

Indications for Use

Section 1.3, page 10

510(k) No.:

Device Name: 3D4-8ET for use with SONOACE R7

Intended Use: Diagnostic ultrasound imaging or fluid flow analysis

	Clinical Application				Mode of (Operation (*inch	des simultaneous B	mode)
General (Track Lonly)	Specific (Tracks I & III)	В	М	PWD	CWD	Color Doppler*	Combined* (Spec.)	Other (Spec.)
Ophthalmic	Ophthalmic							(5500.)
·	Fetal (Nee Note 3)	Р	Р	p		P	No. 1	
	Abdominal	Ρ	P	P		'	Note I Note I	Note 2, 7, 8
	Intra-operative (See Note 6)		_			 - 	Note 1	Note 2, 7, 8
	Intra-operative (Neuro.)					 		
Fetal Imaging	Laparoscopic	1						
& Other	Pediatric	Р	þ	12		P	No. 1	
	Small Organ (See Note 5)	1		-			Note I	Note 2, 7, 8
	Neonatal Cephalic	1	-					
	Adult Cephalic	1-						· — — —
	Trans-rectal	1 -						
	Trans-vaginal	+						
	Trans-crethral	1	-					
	Trans-esoph. (non-Cardiae)	1-						
	Musculo-skel. (Convent.)	┸						
	Musculo-skel. (Superfie.)	╂╌┤						
	Intra-luminal	╂╾┤						
	Other (spec.)	+ +	- 1				————	
	Cardiac Adult	╅						
Cardiac	Cardiac Pediatric	1-						
	Trans-esophageal (Cardiac)	1-	\dashv					
	Other (spec.)	+					·	
Peripheral	Peripheral vessel	+	-+					
Vessel	Other (spec.)	┨─┤						

N= new indication; P= previously cleared by FDA K093714; E= added under Appendix E

Additional Comments:

Color Doppler includes Power (Amplitude) Doppler

Note 1: B/M, B/PWD, B/Color Doppler, B/Color Doppler/PWD, B/Power Doppler/PWD, B/Color Doppler/Color M

Note 2: Includes imaging for guidance of biopsy Note 3: Includes infertility monitoring of follicle development

Note 4: Color M-mode

Note 5: For example: thyroid, parathyroid, breast, scrotum and penis in adult, pediatric and neonatal patients

Note 6: Abdominal organs and peripheral vessel

Note 7: Tissue Harmonic Imaging (THI)

Note 8: 3D imaging

Note 9: Panoramic imaging

Concurrence of CDRH, Office of In Vitro Diagnostic Devices (OIVD) Prescription Use (Per 21 CFR 801.109)

Indications for Use

Section 1.3, page 11